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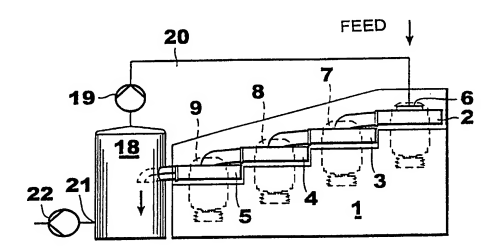
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(54) Title: APPARATUS AND PROCESS FOR THE PREPARATION OF PRECIPITATED CALCIUM CARBONATE



(57) Abstract

The invention concerns a process and an apparatus for the preparation of precipitated calcium carbonate. In the process, calcium hydroxide is carbonated using gaseous carbon dioxide, whereby the carbonation is performed in gaseous phase by mixing calcium-hydroxide-containing liquor mist with the carbon dioxide gas in a turbulence having an energy intensity in excess of 1000 kW/m³. The reaction is advantageously carried out in the apparatus of the invention, comprising at least two serially arranged pin mills (2-5) having one or more rotatable vane rings which can be used to impose a great energy intensity on the material which is fed into the apparatus, whereby the first pin mill is provided with at least an inlet (6) for slaked lime and carbon dioxide and an outlet for the reaction product, and the second pin mill is provided with an inlet (7) for the product from the previous mill and an outlet for the reaction product. According to the invention, carbonating is extremely swift. The residence time of the reaction is even less than 1 second. Due to the high energy intensity, carbonation may be performed at high solids contents.

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Apparatus and process for the preparation of precipitated calcium carbonate

The present invention relates to a process according to the preamble of claim 1 for the preparation of precipitated calcium carbonate (PCC).

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According to such a process calcium hydroxide is carbonated using gaseous carbon dioxide.

The invention also relates to an apparatus according to the preamble of claim 11 for carrying out the carbonation reaction and to a pigment product according to the preamble of claim 15 based on precipitated calcium carbonate.

Precipitated calcium carbonate is used as a paper filling or coating agent. Other possible fields of use include paints, plastics, the food processing industry, the pharmaceutical industry, etc.

PCC can be prepared by a causticizing process and by a carbonation reaction. In the causticizing process the calcium oxide is slaked whereby calcium hydroxide is formed which is then reacted with sodium carbonate in liquid phase. As a result, caustic soda (NaOH) and calcium carbonate are obtained, the sodium hydroxide remaining in dissolved state while the calcium carbonate is precipitated. Both products are recovered and forwarded to further processing.

In the carbonation process, the calcium hydroxide slurry obtained from slaking lime is reacted with gaseous carbon dioxide. This is typically carried out by charging a carbon-dioxide containing gas, derived from flue gas and having a CO₂ content of about 20 to 40 %, into the Ca(OH)₂ mixture whose solids content is about 20 %. The CO₂ gas is hereby blown into an aqueous Ca(OH)₂ solution whereby the gas is disintegrated into bubbles and the carbon dioxide contained in these bubbles is

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dissolved in the surrounding water. Carbonate ions are formed which react with the Ca²⁺ ions whereby calcium carbonate is obtained which is precipitated from the solution.

In order to provide an exhaustive account it may be mentioned that calcium hydroxide is reacted with carbon dioxide not only in the preparation of PCC but also in other contexts such as, among others, desulphuration of flue gases and scrubbing in a flue gas scrubber.

A number of considerable disadvantages are related to the prior art processes for preparing PCC. Thus, the conventional causticizing process is hampered by residual salts in the PCC. When, on the other hand, PCC is produced by conventional carbon dioxide carbonation processes, a disadvantage lies in the long carbonation time, typically 1 to 7 h, required by the reaction. In addition, the PCC crystals produces are of varying size and their particle size varies within a very wide range.

The prior art solutions have also been hampered by the difficulty of attaining sufficiently efficient mass transport conditions to enable fast nucleation and the simultaneous generation of a vast number of crystal seeds which would then grow into a vast number of small crystals.

The aim of the present invention is to remove the drawbacks of the prior art and to obtain an entirely novel solution for the preparation of precipitated calcium carbonate from slaked lime and carbon dioxide gas.

The invention is based on the concept of performing the carbonation subject to strong turbulence in a turbulence zone by reacting the carbon dioxide gas with calcium hydroxide particles by the intermediation of random liquid droplets. Thus, in the reaction, gas, liquid and solids particles are contacted with each other simultaneously under intense turbulence and a great energy intensity. The gas flow absorbs the liquid and the particles and forms a turbulent three-phase mixture. The solution according to

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the invention can also be termed a three-phase process because three phases are simultaneously present, the gaseous phase constituting the reaction medium.

In more detail the process according to the invention is mainly characterized by what is stated in the characterizing part of claim 1.

The apparatus according to the invention comprises at least two serially arranged pin mills having one or more rotatable vane rings by means of which it is possible to subject the material charged into the apparatus to a great energy intensity. The first pin mill is furnished at least with an inlet for slaked lime and carbon dioxide and a discharge outlet for the reaction product and the second pin mill is furnished with an inlet for the product from the previous pin mill and a discharge outlet for the reaction product. Gas or blend liquor can, if desired, be fed between the rotating vane rings or groups of vane rings of the pin mills. The pin mills are connected to each other by means of pipes which can be furnished with inlets for blend liquors, if-desired.

In more detail the apparatus according to the invention is characterized by what is stated in the characterizing part of claim 11.

By means of the invention, completely novel products can be obtained having a shell-like structure. The precipitated particles preferably have a size of 30 to 100 nm and are of spherical shape whereby they are formed of one or several shell-like layers.

The product is characterized by what is stated in the characterizing part of claim 15.

The invention offers considerable benefits. Thus, the carbonation of calcium hydroxide is extremely swift. The dwell time of the reaction may be as short as less than 1 second. Due to the great energy intensity carbonation may be performed at a high solids content (even at 40 to 60 % by w.)

The calcium carbonate obtained by the invention is of homogeneous quality; the

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diameters of the PCC produced may be e.g. 20 to 30 nm, 30 to 50 nm and 50 to 100 nm, i.e. generally within the range from 20 to 100 nm, usually 30 to 100 nm. The minute PCC particles produced can be exploited in various ways: by combining them to form bigger pigment particles by means of van der Waals forces, particle clusters are obtained containing 10 to 30, typically about 15 to 20 joined particles. The formation of these particle clusters can be carried out by adjusting the pH to a value within the range from 6.2 to 10.8 whereby the Z potential of the particles is as small as possible. The particles can also be used for coating other pigments such as kaolin, chalk, talc, or titanium dioxide. The coating can be carried out by feeding the pigments to be coated e.g. in the form of an aqueous slurry together with calcium hydroxide and carbon dioxide into the apparatus of the invention and, if needed, by adjusting the pH value to a suitable range e.g. by introducing acid into the pin mill apparatus during production.

The carbonation is divided into several (e.g. 3 to 7) different process stages. The conversion of calcium carbonate increases step by step; depending on the dry matter content of the calcium carbonate it is usually close to 100 after 3 or 4 stages already. By dividing the process into stages blend components can be added to the different layers of the CaCO₃ particle, said components affecting, among other things, the opacity and acidity resistance of the product. As an example a product may be cited prepared by a multi-step process wherein the obtained particles have a core layer consisting of calcium carbonate and a few shell layers consisting alternately of calcium phosphate and calcium carbonate, and a surface layer consisting of, e.g. calcium phosphate. Such a structure will improve the acid resistance of the calcium carbonate particles. In addition, variations in the refractive index between the different layers will provide improved opacity as compared to a mere CaCO₃ particle.

In the following, the invention is examined in more detail by means of a detailed description, the annexed drawings and a number of working examples.

Figures 1a and 1b provide a simplified side view and correspondingly a top view of

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the principal structure of an apparatus consisting of four serially arranged pin mills.

Fig. 2 is a sectional side projection of a pin mill and

Figs 3 and 4 are sectional top views of single and correspondingly double rotor mills.

It has been found according to the invention that mass transport conditions in a carbonation reaction can be made highly efficient in gaseous phase. Because the density of gas is smaller than that of liquid, a mixing intensity is achieved in gaseous phase which only requires about 1/1000 of the energy which would have to be used in liquid phase to achieve an equivalent mixing intensity. Subject to turbulence, the gas is reacted with solid particles being in the same mixing state by intermediation of liquid droplets (i.e., almost water droplets).

The carbonating process of the present invention is performed in an aerosol phase, i.e. gaseous phase, water droplets having been dispersed therein and containing mainly $Ca(OH)_2$ as reagent. When a mist of this type is made to repeatedly impinge on the actuator providing the kinetic energy or a gas turbulence generated by the actuator in a rapidly rotated flow channel, the surface is continuously renewed providing a high nucleation rate and, in the end, a great number of minute particles.

According to the invention CO₂ gas is therefore subjected to a strong turbulence having an energy intensity of > 1,000 kW/m³. Ca(OH)₂ slurry is fed into this state, the solution having a solids content of < 70 %, preferably between 5 and 50 %. The volume fraction of the Ca(OH)₂ solution/slurry of the gas volume of the apparatus is small, typically smaller than 1 %, preferably about 0.1 to 5 ‰. To cite an example, an apparatus having a gas volume of about 40,000 cm³ may be charged with approximately 10 to 200 cm³, advantageously about 50 to 150 cm³, of a calcium hydroxide slurry, and an energy of approximately 2,000 kW/m² is impinged on this aerosol.

In the turbulence the water droplets are converted into mist and their surface area increases, whereby the CO₂ is rapidly dissolved in water. The mist-like water and the

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water in the diffusion layer on the particle surface stand in efficient interaction. Therefore, the small size of the water droplets produces a large contact surface area and accelerates dissolution. The particles collide into each other producing temperature peaks which for their part accelerate the reaction. Thus, after four turbulence stages a 100 % carbonation has taken place with a 20 % CaCO₃ solution.

The reactions of reactants absorbed in water are converse, i.e. they take place in both directions depending on what forms of occurrence are consumed in the reaction.

Any apparatus is used for providing the turbulence, i.e. as the turbulence zone, capable of producing a high energy intensity in the gas volume. Advantageously, the apparatus is a so called pin mill or a corresponding device (shock mixer) or a bead mill. An advantageous apparatus is described in, e.g. WO Published Application 96/23728. As a rule, the apparatus in question is filled with reagent gas and only contains small volumes of materials in, e.g. liquid or solid phase. This condition can also be met in, e.g. a disc or cone refiner, which are constructed for a totally different purpose.

The turbulence can be generated in one or more apparatuses. It is of particular advantage to perform the reaction in several serially (successively) arranged mixers whereby the same continuous renewal of the surface of the reagent film is performed again and again.

According to the invention it has been found that particularly good results are achieved if carbon dioxide gas of maximum purity is introduced into the reaction. The purity of the CO_2 gas should preferably exceed 90 %. Correspondingly, advantageous results are obtained with $Ca(OH)_2$ particles having a size which is < 1 μ m \emptyset . The water used should contain little or no Fe, Mn, or other metals.

Figs 1a and 1b illustrate an apparatus according to the invention where four pin mills 2 - 5 are arranged in series on a steady frame 1. The pin mills may comprise, e.g.

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single and/or double rotor mixers. Thus, the apparatus comprises vane rings rotating in different directions, or a rotating vane ring and a non-rotating vane ring. A pair of rotors or a pair of a rotor and a stator can be provided with, e.g., five vane rings. The discharge pipe 10 - 13 from each pin mill 2 - 4 is connected to the inlet 7 - 9 of the next pin mill. The feedstock supply, i.e. the supply of slaked lime and carbon dioxide gas occurs through the inlet 6 of the first pin mill. The obtained fluid is transmitted from the outer periphery to the next mixer 3 due to centrifugal forces and underpressure, and from there it is transmitted further to the following mixers 4, 5. The pin mills are driven by actuators 14 - 17.

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The discharge pipe 13 from the last pin mill 5 is fitted inside a gas separator tank or a pumping tank 18. In the pumping tank 18 the fluid is separated into a CaCO₃ mixture and a gas which mainly comprises CO₂ and aqueous vapour. The CO₂ gas is returned to the first mixer 6 of the arrangement via the pump 18 and the recirculation line 20 in order to be reutilized in the process. The product is removed from the tank through the discharge outlet 21 using the pump 22. The CaCO₃ mixture can be used as pigment either as such or after a finishing treatment.

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The benefit of the arrangement is that mixture components can be fed into the turbulence at the different intermediate stages of the carbonating process. Thus, more CO_2 gas and mixture components can be fed into the connecting pipes between the mixers (i.e. the discharge pipes 10 - 13 of the pin mills).

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The apparatus can also be arranged as one device such that a multi-periphery rotor is constructed having a diameter in accordance with the example and the mixture components are fed into the mixing chamber at the stators.

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Figs 2 to 4 provide sectional side and, correspondingly, top views of the pin mill used in the invention. The pin mill has a drum 31 of fairly low height and a feed orifice (inlet) 32 is provided in the upper part thereof. One or more vane rings, or grinding peripheries 33, 34, are arranged inside the drum such that at least one of the rings is

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rotatably mounted on bearings. The second grinding periphery is statically mounted or rotatable. The planar circumferential disks of the grinding rings are equipped with perpendicular pins 35. In Fig. 3 a double-ring pin mill is shown wherein both grinding rings are rotatable, and Fig. 4 illustrates an embodiment where stators with perpendicular grinding pins are provided between the rotatable pin rings.

As will emerge from Fig. 2, the gap between the sets of grinding rings can be arranged such that it expands in the radial direction.

In addition, a tangential discharge pipe 36 is fitted to the grinding drum.

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The densely dashed lines indicate the path of the solid matter/liquid being processed through the pin mill.

As stated above, special benefits are gained by a serial arrangement of several turbulence zones. These can, however, also be replaced by one single pin mill. Thus, an arrangement corresponding to a triple mixer combination can be achieved by a 1400 ø rotor/stator combination having 11 to 15 rotating vane rings. Alternatively, one double-rotor mixer with 5 rings can be combined with one single-rotor mixer with 10 rings. In such a combination it is the double-rotor mixer that prepares the fluid, and the single-rotor mixer processes the fluid further.

The apparatus according to the invention can be used for the preparation of calcium carbonate as well as for the modification of calcium carbonate and other pigments. In the latter case, pigments can, for example, be coated with PCC particles which are used to improve the optical properties of the pigments. According to an advantageous embodiment, blend components are introduced into the carbonating reaction or the modification of pigments. Examples of suitable blend components include (NaPO₃)₆, phosphoric acid, hexameta-, pyro-, tripoly-, poly- or ultraphosphoric acid, aluminium T, silicic acid chloride or fluoride of aluminium, aluminium sulphate. The blend components can be charged into the reactor in gaseous form.

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As an example, an embodiment can be cited wherein the aim is to improve the acid resistance of calcium carbonate. Hereby phosphoric acid H₃PO₄ (or a phosphoric acid derivative) is fed into the apparatus in addition to carbon dioxide, and the phosphoric acid is gasified. As solid matter, calcium hydroxide can be used, which is carbonated at the same time, or calcium carbonate prepared earlier with the apparatus, the calcium carbonate being then coated in the apparatus by feeding it again through the apparatus together with the blend components. Both ways are applicable to the treatment of other pigments, too.

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First alterative:

H₃PO₄

gas

 H_2O

water

CaCO₃

particles

Second alternative: CO₂ gas

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 $Ca(OH)_2 + H_2O$ mixture

powder or mixture of powder + water

- kaolin
- titanium dioxide
- lime (CaCO₃)

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- ground limestone
- CaCO₃ (precipitated CaCO₃ (PCC))

An additive is added to the above during an intermediate stage in the process, whereby products differing from CaCO₃ as regards their opacity and acid resistance are obtained.

The obtained particles contain e.g. the following:

Core

CaCO₃

Layer

 $Ca_3(PO_4)_2$

Layer

CaCO₃

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Surface $Ca_3(PO_4)_2$

Variations in the refractive index between the different layers provide improved opacity as compared to a mere CaCO₃ particle.

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The following examples are provided by way of illustrating the invention without limiting its scope of protection. The examples were implemented in the apparatus of Fig. 1 with the following components generally present in the turbulence volume:

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gas	40,000 cm ³
liquid	80 cm ³
particles	20 cm ³

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Example 1

Reaction Ca(OH)₂ + CO₂ - CaCO₃ + H₂O

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Test apparatus Single-rotor mixer energy intensity 2,000 kW/m³

Gas Twice the equivalent amount of CO₂ - 100 %

Ca(OH)₂ mixture solid matter content 5 %

water 95 %

Table 1

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Result	,	Time s	Temperature °C CaCO ₃ %		
		Start	53	< 10	
Stage	I ≈	0.04	42	66	
Stage		0.04	42	97	
Stage		0.04	38	100	
Stage		0.04	35	100	
J	<	1 s			

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Example 2

Seaction $Ca(OH)_2 + CO_2 - CaCO_3 + H_2O$

Test apparatus

Single-rotor mixer, energy intensity 2,000 kW/m³

Gas

 CO_2 - 100 % feed 2 x the equivalent amount

Ca(OH)₂ mixture

solid matter content 10 %, water 90 %

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Table 2

		Time s	Temperature °C CaCO ₃ ,%		
		-	40	210	
I		0.04	46	59	
II		0.04	44	90	
III		0.04	43	99	
IV		0.04	40	100	
	<	1 s			

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Example 3

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Reaction Ca(OH)₂ + CO₂ - CaCO₃ + H₂O

Test apparatus

Double-rotor mixer, energy intensity 3,700 kW/m³

Gas

CO₂ - 100 % feed 2 x the equivalent amount

Ca(OH)₂ mixture

solid matter content 20 %, water 80 %

Table 3

	Time s	Temperature °C CaCO ₃ %		
		40	< 10	
T	0.04	46	. 41	
II	0.04	47	76	
III	0.04	47	97	
IV	0.04	47	100	

Example 4

Reaction $Ca(OH)_2 + CO_2 - CaCO_3 + H_2O$

Test apparatus Double-rotor mixer, energy intensity 5,500 kW/m³

Gas CO₂ - 100 % feed 2 x the equivalent amount

Ca(OH)₂ mixture solid matter content 50 %, water 50 %

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Table 4

			Time s	re °C CaCO ₃ %	
25			•	50	< 10
	I		0.04	45	35
	II		0.04	44	64
	III		0.04	44	86
	IV		0.04	44	98
30	V		0.04	44	100
		<	l s		

35 Example 5

Reaction $Ca(OH)_2 + CO_2 - CaCO_3 + H_2O$

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Test apparatus	Double-rotor mixer, energy intensity 3,700 kW/m ³
Gas	CO ₂ - 25 %, feed 2 x the equivalent amount, air - 75 %
Ca(OH), mixture	solid matter content 10 %, water 90 %

5 Table 5

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		Time s	Temperature °C CaCO ₃ %		
		•	33	< 10	
Ţ		0.04	38	25	
ÎI	æ	0.04	38	38	
III	æ	0.04	38	48	
IV	æ	<u>0.04</u>	39	62	
	>	1 s			

As will emerge from the above table, air admixed with the CO_2 hampers the reaction of CO_2 with the $Ca(OH)_2$ particle.

Claims:

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- 1. A process for the preparation of precipitated calcium carbonate, wherein calcium hydroxide is carbonated with gaseous carbon dioxide, **characterized** in that the carbonating is performed in gaseous phase by contacting calcium hydroxide and liquid mist with carbon dioxide gas in a turbulence having an energy intensity in excess of 1,000 kW/M³.
- 2. The process according to claim 1, characterized in that the volume of the liquid is less than 1 % of the volume of the gas.
 - 3. The process according to claim 1, characterized in that the carbon dioxide has a degree of purity of at least 90 %.
 - 4. The process according to any one of the previous claims, characterized in that the carbon dioxide and the calcium hydroxide are subjected to an energy intensity of approximately 1,100 to 6,000 kW/m³.
 - 5. The process according to any one of the previous claims, **characterized** in that the calcium hydroxide is carbonated in several serially arranged turbulence zones.
 - 6. The process according to claim 5, characterized in that pigments which are coated with the calcium carbonate produced are fed into at least one turbulence zone.
 - 7. The process according to claim 6, **characterized** in that kaolin, titanium dioxide, chalk, ground limestone or precipitated calcium carbonate is fed into at least one turbulence zone.
 - 8. The process according to any one of the previous claims, characterized in that the carbonating is performed in a turbulence zone comprising a pin mill.

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9. The process according to any one of the previous claims, **characterized** in that reactive gas is fed into the turbulence zone in order to modify the calcium carbonate, the gas comprising hydrogen fluoride, aluminium chloride, silicic acid chloride or fluoride, or titanium tetrachloride.

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10. The process according to any one of the previous claims, **characterized** in that the gas-transmitting liquid droplets are in atomized form.

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11. An apparatus for the preparation of calcium carbonate by carbonating calcium hydroxide, whereby carbon dioxide is reacted with calcium hydroxide via liquid mist, characterized in that the apparatus comprises

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at least two serially arranged pin mills having one or more rotatable vane rings which can be used to impose a high energy intensity on the material introduced into the apparatus,
a first pin mill which is equipped with at least an inlet for slaked lime and

carbon dioxide and an outlet for the reaction product,
- a second pin mill which is equipped with an inlet for the product of the
first pin mill and an outlet for the reaction product, and

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- pipes interconnecting the pin mills, which pipes can be provided with inlets for the blend liquors, if desired.

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12. The apparatus according to claim 11, **characterized** in that the pin mills are provided with inlets which can be used to feed gas or blend liquor between revolving vane rings or groups of vane rings.

13. The apparatus according to claim 12 or 13, characterized in that it comprises 3 to 7 serially arranged pin mills.

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14. The apparatus according to any one of the claims 11 to 13, characterized in that the pin mill is furnished with a double-rotor and/or a single-rotor-stator mixer or mixer group.

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- 15. A pigment product based on precipitated calcium carbonate, characterized in that the precipitated particles are spherical in shape and have a size of 30 to 100 nm and are formed of one or several shell-like layers.
- 16. The pigment product according to claim 15, **characterized** in that the precipitated particles contain a core layer consisting of calcium carbonate, a few shell layers consisting alternately of calcium phosphate and calcium carbonate and a surface layer of calcium phosphate.

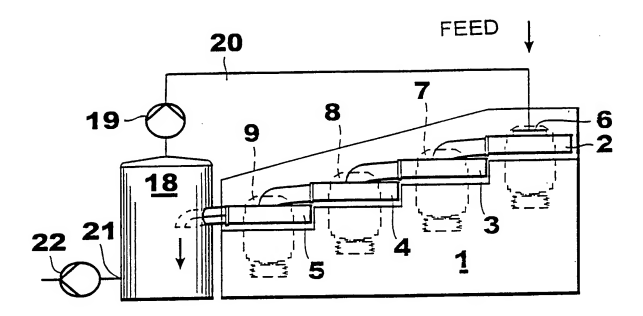
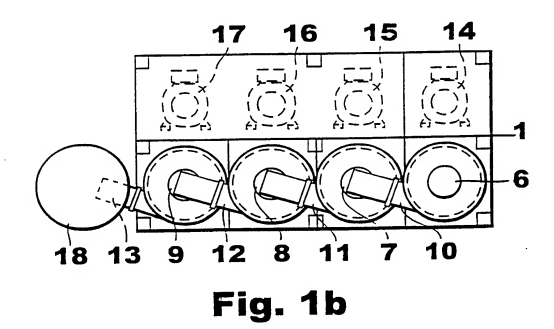
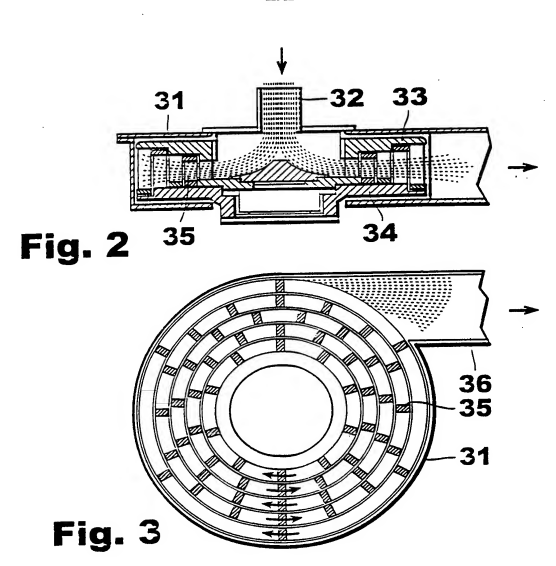


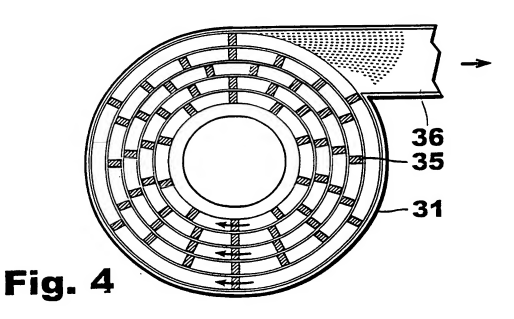
Fig. 1a



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International application No.

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A. CLASS	IFICATION OF SUBJECT MATTER								
IPC6: C01F 11/18 According to International Patent Classification (IPC) or to both national classification and IPC									
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Minimum do	ocumentation searched (classification system followed by	classification symbols)	. 						
	Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched								
	SE,DK,FI,NO classes as above								
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WPI									
C. DOCU	MENTS CONSIDERED TO BE RELEVANT								
Category*	Citation of document, with indication, where app	ropriate, of the relevant passages	Relevant to claim No.						
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Furth	ner documents are listed in the continuation of Box	C. X See patent family anne	x.						
"A" docum	l categories of cited documents: tent defining the general state of the art which is not considered of particular relevance	"T" later document published after the int date and not in conflict with the appl the principle or theory underlying the	ication but cited to understand						
"E" erher o	document but published on or after the international filing date tent which may throw doubts on priority claim(s) or which is o establish the publication date of another citation or other	"X" document of particular relevance: the considered novel or cannot be considered novel or cannot be considered when the document is taken along the considered to the consid	ered to involve an inventive						
"O" docum means	reason (as specified) tent referring to an oral disclosure, use, exhibition or other	"Y" document of particular relevance: the considered to involve an inventve ste combined with one or more other such being obvious to a person skilled in the consideration of th	p when the document is h documents, such combination						
	ent published prior to the international filing date but later than ority date claimed	"&" document member of the same patent							
Date of th	ne actual completion of the international search	Date of mailing of the international	search report						
7 July	1998	09-07-	1998						
Name and	d mailing address of the ISA?	Authorized officer							
Box 5055	Patent Office 5, S-102 42 STOCKHOLM No. + 46 8 666 02 86	Britt-Marie Lundell Telephone No. +46 8 782 25 00							
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International application No.
PCT/F198/00244

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This inte	mational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2.	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	mational Searching Authority found multiple inventions in this international application, as follows:
	See extra sheet
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-14.
Remark	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

International application No. PCT/F198/00244

Invention 1, claims 1-14, relates to a method and a device for preparing precipitated calcium carbonate by carbonating calcium hydroxide with gaseous carbon dioxide. The invention is characterised by reacting the components under high turbulence.

Invention 2, claims 15-16 relates to a pigment product, characterised by its physical form and size.

The inventions 1 and 2 are not so linked as to form a single general inventive step.

The special technical features of invention 2 is not corresponding to the technical features of invention 1.

Information on patent family members

International application No.
PCT/FI 98/00244

	atent document d in search repor	t ,	Publication date	_	Patent family member(s)		Publication date	
US	4888160	A	19/12/89	US US	5055284 5338777		08/10/91 16/08/94	
WO	9206038	A1	16/04/92	AT AU CA DE EP SE ES FI GB JP US	649721 8526291 2087857 69105183 0480587 0480587 2064047 930143	A A D, T A, B T3 T A A, B	15/12/94 02/06/94 28/04/92 28/03/92 24/05/95 15/04/92 16/01/95 14/01/93 01/04/92 10/02/94 30/08/94	
WO	9623728	A1	08/08/96	AU FI FI FI	4541696 2182 100237 950411	U B	21/08/96 17/11/95 00/00/00 31/07/96	